

ERDC/CRREL TR-03-22

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A Preliminary Investigation of Diffusion of Volatile Atmospheric Acids in Snow

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November 2003

20040329 080

ABSTRACT

A preliminary coldroom experiment was conducted to determine the diffusion rate of several reactive atmospheric chemical contaminants in snow at -7°C . These species included formaldehyde, p-nitrotoluene (PNT), HNO_3 , and HCl . Because of detection or contamination problems we could not quantify diffusion rate for formaldehyde, PNT, or HNO_3 . However, the diffusivity of HCl in snow ($\rho = 0.38\text{ g/cm}^3$) was found to be $> 7.7 \times 10^{-4}\text{ cm}^2/\text{s}$. This is about two orders of magnitude greater than hitherto expected and suggests that HCl may be rapidly redistributed or lost from seasonal snowpacks. Thus, one should be cautious in interpreting temporal records of HCl and other volatile species in seasonal or polar snowpacks, until their integrity is better established.

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PREFACE

This report was prepared by James H. Cragin, Research Chemist, Snow and Ice Branch, U.S. Army Engineer Research and Development Center (ERDC), Cold Regions Research and Engineering Laboratory (CRREL), Hanover, New Hampshire; and by Daniel C. Leggett, Research Chemist, Environmental Sciences Branch, ERDC-CRREL.

Funding was provided by ERDC/CRREL.

The authors thank Dr. Samuel Colbeck and Dr. Matthew Sturm of CRREL for technically reviewing the manuscript.

The Commander of the Engineer Research and Development Center is Colonel James R. Rowan, EN. The Director is Dr. James R. Houston.

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JAMES H. CRAGIN AND DANIEL C. LEGGETT

1 INTRODUCTION

Trace constituents of the atmosphere can contribute to important processes such as ozone destruction, acid precipitation, and global warming. It is important to understand the sources and fates of these compounds in order to assess the role of man's activities on these processes. Some trace species that have been implicated are the mineral acids hydrochloric, sulfuric, nitric, and low molecular weight oxygenated species, formate, acetate, formaldehyde, sulfur dioxide, and hydrogen peroxide. These species are very soluble in water and are efficiently scavenged and dissolved by small water droplets in the atmosphere. When these droplets are quickly frozen, as often occurs in the atmosphere, the trace species are "frozen in" (Iribarne and Pyshnov 1990) at concentrations exceeding their equilibrium solubility in ice.

Permanently frozen precipitation may then serve as a semi-permanent record of historic atmospheres, and polar ice core analysis could be used retrospectively to establish past conditions. One may doubt this conjecture, however, as volatile species would be expected to redistribute over time so as to achieve local ice/air equilibria. We would expect these changes to be reflected in temporal changes in volatile species concentrations in snowpacks, and, in fact, these kinds of changes have been observed in mid-latitude snowpacks (Hogan et al. 1985) and in laboratory-stored snow (Cragin and McGilvary 1995). The kinetics of redistribution processes and the ultimate fate of various contaminants remain obscure, however.

Contaminants also enter snowpacks by dry deposition on the fallen snow. This applies not only to atmospheric sources but to sources such as soil beneath the snow. Because of the prevailing temperature gradients, volatile contaminants tend to accumulate in the coldest snow, which is often the top of the pack (Hogan et al. 1985, Hogan and Leggett 1995). Because these phenomena are hard to study in nature, most of our information comes from laboratory investigations.

However, no direct information on diffusion of trace volatile species in snow was found. Diffusion of solute vapors through a porous medium such as snow is expected to be retarded with respect to vapor movement in air by interaction with the media and tortuosity effects.

Available information suggests that interaction of acidic species with ice particles is strong, and that diffusion into the bulk ice occurs. The diffusivities of hydrochloric and nitric acids from dilute vapor sources in ice have recently been estimated. The reported values range from about 10^{-12} cm²/s at -8° C for ice single crystals (Domine and Thibert 1995) to around 10^{-8} cm²/s for polycrystalline ice (Diehl et al. 1995). The latter value would presumably be more relevant for atmospheric studies and for the work reported here since snow is essentially unconsolidated polycrystalline ice. If so, we would anticipate essentially complete permeation of the ice spheres during the time course of our experiment, although complete equilibration with the ice lattice could take years, assuming the lower diffusivity.

To avoid complicating effects of temperature gradients, we used a single isothermal temperature for this preliminary study of dilute vapor migration through snow. Compounds initially selected for study were HCl, HNO₃, formaldehyde, and p-nitrotoluene. The latter was added as a reference compound as one of us had successfully used it before in outdoor, temperature gradient experiments (Hogan and Leggett 1995).

2 EXPERIMENTAL METHODS

All laboratory work was performed in a Class 100 cleanroom or Class 100 clean air station in a coldroom. Because of the finite background concentrations of many atmospheric species in natural snow, we prepared clean artificial snow from frozen Milli-Q deionized water as described below.

First, we added Milli-Q water to pre-cleaned 7.5-cm-diameter polyethylene tubes that were heat-sealed on the bottom. Leaving about 20 cm of air space for expansion above the 60 cm of water, we sealed the top and slowly (1.8 mm/hr) froze the water vertically by lowering it into a circular opening in a freezer (see Cragin and McGilvary [1995] for a more detailed description of the apparatus and process). We sliced the resulting ice core into 1-cm-thick sections and rinsed them in Milli-Q water to remove surface contaminants and then mechanically crushed the sections in a pre-cleaned Plexiglas chamber (similar to a mortar and pestle).

We sieved the resulting snow so that the 0.425- to 0.850-mm-size range fell naturally through a funnel into a 10-cm-diameter Plexiglas column. We suspended separate 1-mL vials of formaldehyde, 0.1 M HCl, 0.1 M HNO₃, and PNT in the airspace above the snow and finally sealed the column with an airtight cap. Vapor was free to move within the chamber from the uncapped source vials to the snow. We left the apparatus in a coldroom at -7° C.

After one week we opened the column, removed the source vials, and sectioned the snow core crosswise in a clean air station in the coldroom. The resulting 1- or 2-cm-thick wafers were melted and analyzed. Analyses for formaldehyde and PNT were done using gas chromatography while chloride and nitrate were determined using ion chromatography.

3 RESULTS AND DISCUSSION

We failed to detect p-nitrotoluene (PNT) in any of the snow samples. This was somewhat surprising because it had been detected in earlier work (Hogan and Leggett 1995) in snow over PNT-spiked soil at a similar temperature and using the same method of analysis. The difference was that the earlier experiment was conducted outside in the presence of a strong temperature gradient, so that the source was at a higher temperature ($\sim 0^{\circ}\text{C}$) compared to -7°C in the present experiment. The lower temperature is expected to reduce the source strength by a factor of between 2 and 3. The temperature of the snow in which PNT was detected in the former experiment may also have been a bit colder than -7°C , and we failed to detect it when the snow was warmer than -5°C . Evidently, the sensitive temperature dependence of PNT source strength and retention slow its diffusion under isothermal conditions. This seems likely to be a very general phenomenon rather than unique to PNT.

We also did not detect nitrate in any of the snow samples, but this may be because concentrations were below the instrumental detection limit. Because of its high vapor pressure, we expected nitric acid to volatilize and diffuse readily through the snow, but if nitrate concentrations were below $20\text{ }\mu\text{g/L}$, they would have been undetected.

Attempts to determine formaldehyde likewise proved frustrating. The problem was not in failure to detect, however, but in unexpected contamination. Unfortunately, the original ice used to make the snow for the column, which was not analyzed until after the experiment, was contaminated. Similar amounts were also found in all column samples, which precludes drawing any conclusion except that the results are not inconsistent with formaldehyde diffusion through the column. A definitive result must await further experimentation. Analysis of several other batches of lab Milli-Q-purified water also had varying amounts of contamination. One sample of well water from a local source was analyzed and revealed undetectable levels of formaldehyde.

Before future experiments are conducted, potential sources of laboratory contamination other than water, such as exposure to contaminated lab air, must also be ruled out. Indoor air generally contains more formaldehyde than outdoors due to degassing of materials containing formaldehyde in a confined space. Since formaldehyde occurs ubiquitously in the environment it will probably prove impossible to completely eliminate this source of contamination; however, it should be possible to do the experiment by minimizing exposure to sources of contamination and, if necessary, by increasing the formaldehyde source strength.

Figure 1 shows the concentration profile for HCl at the end of seven days. The profile is atypical in that, although the gas has permeated through the entire column, the concentrations behind the front decrease regularly. With a constant source, uniform concentrations behind the front would be expected. The simplest explanation for this result is that the source did not remain constant over time but became depleted. This is prone to happen with unstirred liquid solution sources such as used here, and would be exacerbated by the restricted geometry of the source, because of the mismatch between the rates of gas phase vs. liquid phase diffusion. Diffusion coefficients of solutes are roughly four orders of magnitude higher in air than in aqueous solutions. This can cause depletion of the surface layer of the source and proportionately reduced solute vapor flux when diffusion through the porous medium is as rapid as it was here. Rapid diffusion of HCl through the snow column was not anticipated, as the literature suggested retention by the ice surface would be more efficient (Diehl et al. 1995, Domine and Thibert 1995).

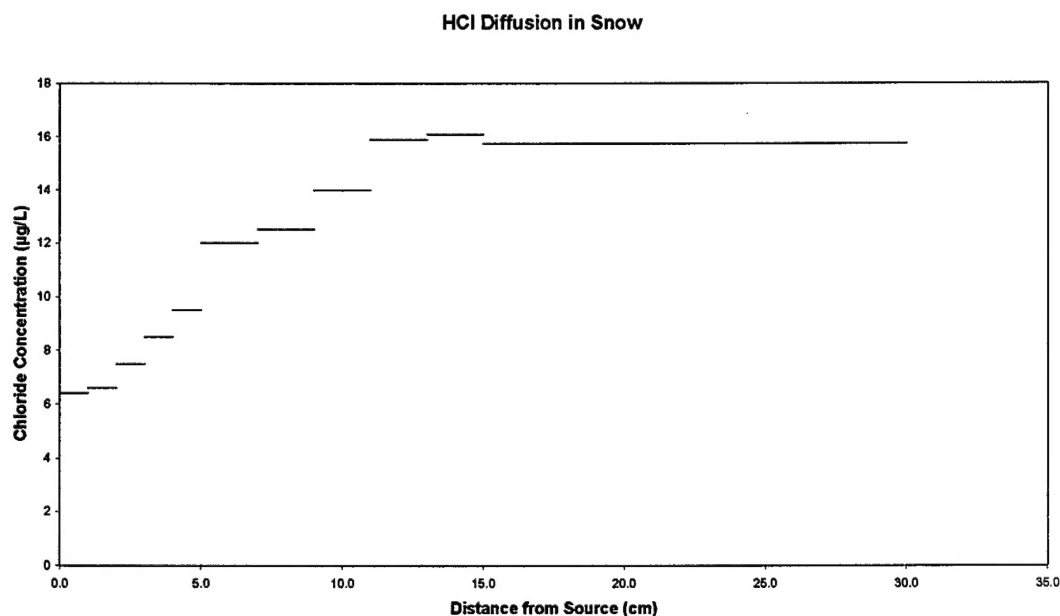


Figure 1. Chloride concentration in snow versus distance from the source (HCl). The total length of the snow column was 30 cm.

One might ask why the HCl doesn't diffuse back through the snow column. It does. Because diffusion occurs in both directions (it actually occurs in three dimensions, but the column wall restricts radial movement), HCl must have

diffused backward out of the snow once the source became depleted. The lower concentrations closer to the source probably reflect this as the HCl diffused out of the snow and possibly repartitioned with solutions in vials of the other sources. Once the HCl became depleted in its source vial, the solutions in the other vials became sinks for HCl in the headspace and the snow.

Since the diffusion front passed completely through the column we cannot calculate an exact diffusivity of HCl for the experiment. We can estimate a minimum value, however, assuming the following equation applies:

$$r = (2Dt)^{1/2} \quad (1)$$

where r is the distance of the front from the source at the end of the experiment, D is the effective diffusivity or diffusion coefficient, and t is time. In this case $r > 30$ cm and $t = 5.83 \times 10^5$ s (6.75 days). Solving for D we obtain a value of $D > 7.7 \times 10^{-4}$ cm²/s. This calculation shows why the HCl flux through the column was limited by mass transfer in the source solution. Diffusivity of HCl in water at -7° C is expected to be no more than 10^{-5} cm²/s (Thibodeaux 1979). Since this is about two orders of magnitude less than its minimum estimated diffusivity in the snow column, it is easy to see why such a mass transfer limitation occurred.

Diffusion of gases in porous media is retarded relative to their diffusion in air by sorption on the media and tortuosity effects. We estimate the effect of tortuosity by assuming spherical geometry of the snow particles and additivity with respect to the two volume fractions, the air-filled voids, and the particle fraction. If we denote these by V_v and ρ_s , the tortuosity factor would be just $V_v + \rho_s (\pi/2)$. The measured bulk density of the snow was 0.38 g/cm³. Therefore an estimate of tortuosity for the experiment is $0.62 + 0.38 (\pi/2) = 1.22$. This is probably a minimum estimate due to the crude assumptions, but seems in reasonable agreement with values reported for other porous media (Thibodeaux 1979).

It was of interest to estimate a sorption coefficient for HCl on snow. In principle the bulk sorption coefficient can be determined from the effective diffusivity and tortuosity. We can't do this since we don't have a definite value for the diffusivity, but we can estimate a maximum value by assuming that $r = 30$ cm, i.e., the length of the snow column. The molecular diffusivity of HCl in air at -7° C was estimated as 0.14 cm²/s from reported values for other compounds of similar molecular weight at 0° C (Thibodeaux 1979; the lower temperature should have negligible effect). We divide this by the tortuosity to obtain a theoretical diffusivity for HCl in the snow column in the absence of adsorption.

Our estimated unretarded diffusivity is then about $0.11 \text{ cm}^2/\text{s}$. An equation for retarded diffusion in porous media is

$$R = D_e/D_r = 1 + (\rho_s K_b/V_v) \quad (2)$$

where D_e is the effective diffusivity without retardation, D_r is the retarded diffusivity, $> 7.7 \times 10^{-4} \text{ cm}^2/\text{s}$, ρ_s is now the bulk density of the snow in g/cm^3 , V_v is the void fraction in mL/cm^3 , and K_b is the bulk sorption coefficient and has dimensions of mL/g . Rearranging in terms of K_b we have

$$K_b = (R - 1) V_v/\rho_s. \quad (3)$$

Substituting the above values gives $K_b < 140 \text{ mL}/\text{g}$.

An alternative expression for sorption coefficient is in terms of surface area. To do this we just divide K_b by the surface area of the snow particles in cm^2/g . We don't have precise knowledge of this but can estimate it from the sieve sizes used to prepare the snow and assuming spherical geometry of the particles. For the sieve openings we used the particles range from 0.425 to 0.850 mm in diameter. This gives a range of specific surface areas of 79 to $158 \text{ cm}^2/\text{g}$. For estimation purposes we use the mean or $120 \text{ cm}^2/\text{g}$ and estimate a surface area-normalized sorption coefficient as $K_s < 1.2 \text{ cm}$ (actually $\text{cm}^3 \text{ air}/\text{cm}^2 \text{ ice}$). This is the more meaningful measure if HCl is adsorbed on the ice surface rather than dissolved in the bulk.

Actually the maximum concentration measured in the current experiment of approximately $16 \text{ ng}/\text{mL}$ represents only $1.3 \times 10^{-6} \text{ g}/\text{m}^2$, which is at most 1% of monolayer coverage. Thus we have no reason to infer sorption into the bulk or postulate uptake into a liquid-like surface layer, in agreement with others (Domine et al. 1995).

Of interest, and as a check on the reasonableness of the above calculation, we can back-calculate the minimum HCl vapor concentration that should have been obtained in the column. Since $K_b = C_b/C_a = 140 \text{ mL}/\text{g}$, where C_b and C_a are the respective HCl concentrations in bulk ice and air, we can solve for C_a . We get $C_a = 1.1 \times 10^{-9} \text{ g}/\text{mL}$ or, in terms of pressure, $5.0 \times 10^{-5} \text{ torr}$. This appears to be too high based on available experimental data and recent fitting models (Miller 1983, Luo et al. 1995). The model supports a partial pressure for HCl in equilibrium with 3.7 wt% HCl in water at -7°C of $\sim 1.0 \times 10^{-6} \text{ torr}$.

However, if the model value is correct, K_b becomes $7.3 \times 10^3 \text{ mL}/\text{g}$ and D_r , $2.5 \times 10^{-5} \text{ cm}^2/\text{s}$. At this diffusivity HCl should have permeated only to about 5

cm in the column instead of all the way through. Resolution of this discrepancy must await future experiments. It is conceivable that the other species, namely HNO_3 and formaldehyde, interfered with sorption of HCl on the snow. We note that the partial pressure of HNO_3 under the experimental conditions (estimated according to Luo et al. 1995) would have been nearly an order of magnitude higher than that of HCl , so it may have competed for sorption sites on the ice surface. We anticipate follow-up experiments in which only one species will be present at a time, and will attempt to measure its actual partial pressure experimentally. This is especially important for these strong acids in water as their activities, i.e., partial pressures, are highly concentration-dependent.

4 ENVIRONMENTAL IMPLICATIONS

Although this work is preliminary, the rapid diffusion of HCl observed in snow at -7°C suggests that this and possibly other mineral acids may be rapidly redistributed and lost from seasonal snowpacks. This might also explain why some acid anions are lost on storage of snow samples (Cragin and McGilvary 1995). In nature, temperature gradients tend to drive contaminants upward through snowpacks (Hogan and Leggett 1995) where concentration gradients can return them to the atmosphere by diffusion and convection. Thus, one should be cautious in interpreting measurements of potentially volatile species in seasonal snows or in polar firn, until their integrity is better established.

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REPORT DOCUMENTATION PAGE				<i>Form Approved</i> OMB No. 0704-0188	
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1. REPORT DATE (DD-MM-YY) November 2003		2. REPORT TYPE Technical Report		3. DATES COVERED (From - To)	
4. TITLE AND SUBTITLE A Preliminary Investigation of Diffusion of Volatile Atmospheric Acids in Snow				5a. CONTRACT NUMBER	
				5b. GRANT NUMBER	
				5c. PROGRAM ELEMENT NUMBER	
6. AUTHOR(S) James H. Cragin and Daniel C. Leggett				5d. PROJECT NUMBER	
				5e. TASK NUMBER	
				5f. WORK UNIT NUMBER	
7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES) U.S. Army Engineer Research and Development Center Cold Regions Research and Engineering Laboratory 72 Lyme Road Hanover, NH 03755-1290				8. PERFORMING ORGANIZATION REPORT ERDC/CRREL TR-03-22	
9. SPONSORING/MONITORING AGENCY NAME(S) AND ADDRESS(ES) U.S. Army Corp of Engineers Washington, DC 20314-1000				10. SPONSOR / MONITOR'S ACRONYM(S)	
				11. SPONSOR / MONITOR'S REPORT NUMBER(S)	
12. DISTRIBUTION / AVAILABILITY STATEMENT Approved for public release; distribution is unlimited. Available from NTIS, Springfield, Virginia 22161.					
13. SUPPLEMENTARY NOTES					
14. ABSTRACT A preliminary coldroom experiment was conducted to determine the diffusion rate of several reactive atmospheric chemical contaminants in snow at -7° C. These species included formaldehyde, p-nitrotoluene (PNT), HNO ₃ , and HCl. Because of detection or contamination problems we could not quantify diffusion rate for formaldehyde, PNT, or HNO ₃ . However, the diffusivity of HCl in snow ($\rho = 0.38 \text{ g/cm}^3$) was found to be $> 7.7 \times 10^{-4} \text{ cm}^2/\text{s}$. This is about two orders of magnitude greater than hitherto expected and suggests that HCl may be rapidly redistributed or lost from seasonal snowpacks. Thus, one should be cautious in interpreting temporal records of HCl and other volatile species in seasonal or polar snowpacks, until their integrity is better established.					
15. SUBJECT TERMS <div style="display: flex; justify-content: space-between; margin-top: 5px;"> Diffusion Ice core records </div> <div style="display: flex; justify-content: space-between; margin-top: 5px;"> Hydrochloric acid Snow </div>					
16. SECURITY CLASSIFICATION OF:			17. LIMITATION OF OF ABSTRACT	18. NUMBER OF PAGES	19a. NAME OF RESPONSIBLE PERSON
a. REPORT	b. ABSTRACT	c. THIS PAGE			19b. TELEPHONE NUMBER (include area code)
U	U	U	U	16	